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Result Page

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Bonding agents on acrylate resin basis and their use to anchorings reactive resin masses as sticking and bonding agents on the basis of insatiated polyester resins or of PU and/or of their education partners are for a long time known. Regularly it concerns two-component systems, whereby one.

Component reactive resin and/or some reaction partner and the other component the hardener and/or, the others c RTI ID=1.15 Ph. AC-/ HTIS more tionspartner contains. Others, usual constituents, like Füllmittell, accelerator, stabilisers, solvents including reactive solvents, dyes and such a thing can be into and/or other component a contained. By mixing the two components then the reaction under formation of a curved product becomes on brought. To the lixture of mooring rings in Boreholes generally cartridges used with two chambers to the separation of the components contained in the cartridge. The cartridge will will become into the Borehole used and by revolving bringing of the mooring ring into a component contained in the cartridge. The cartridge will will become into the Borehole used and by revolving bringing of the mooring ring into its able to make straddling pressure free the load introduction possible into the ground. Before all things for heavy load anchorages such shear connectors in the past preserved have themselves.

Meanwhile adhesive, insatiated polyester resin systems e.g. point to basis from < RTI ID-2.1.5 o-</RTI. and/or isophthalic acid, mark in and/or fumaric acid as dicarboxylic acid and Dialkholen with the risen requirements at adhesion strength and/or to the attachments certain weakness up. Like that these systems borders are in the thermostability (WFB after Martens 50 - 700C) as well as in the resistance to aging, in particular with influence of humidity, temperature changes and alkality set.

PU systems on the other hand react to sensitive with secondary reactions in presence of humidity. Beside the normally running reaction between polycle and iscoryantae a competition reaction with the water occurs. Since iscoryante reaction with higher rate runs off the waters, in addition a portion of iscoryanate under C02-Bildung for the PU formation is lost in uncontrolled way, troubles are in the structure the inevitable succession.

Object of the present invention is it to make adhesive means available which free is from the known drawbacks and beyond that makes further advantages accessible, how they are following in exemplary way discussed.

Inventive one becomes this achieved with radical hardenable sticking, detention and sealant on acrylate basis, those inventive bisphenol and or Novolak connections available reaction products contained exhibiting as hardenable acrylate by conversion of acrylic acid derivatives with Epoxigruppen.

The invention is the basis the realization that by the aforementined conversion available moments or oligomeren, reactive connections particular advantages make accessible. Know mone, particular however dispositions of disponsible or disponsibl

Prefered ones are here derivatives of bisphenol A or F and of Novolak connections, in particular with a content of group of benzyles from \$0.6. As acrylic acid derivatives connections can be used, some free < RTI D=3.1> carbonic acid < /RTI> group or, a functional carboxylic acid derivative contained reactive with Epoxiden. Prefered one is acrylic acid and Methacrylsturederivate and acid derivatives of these two connections like e.g., esters, amides or halides, in particular the chlorides and the derivatives of such connections like e.g., - halogeneous or others - the lowalikyl substituted derivatives. The esters are preferably the esters with alighatic, especially < RTI ID=3.2> CI-CS-alightatischen < RTI SL Alcohols in addition, with henoils and cyclohoxanols.

Particularly preserved have itself reaction products, which by conversion of acrylic acid and/or their derivatives with such bisphenols, which are 2 Epoxigruppen and/or such novolacs, which have a Epoxigruppe per phenol core, available.

Preferably the Epoxireste is connected with the bisphenols and/or novolacs by an ether connection, even it direct CC linkage is not excluded. The ether bridge attacks the Epoxiverbindungen at the phenol core, dh is in particular directly connected under Verätherung of the phenolic OH-groups with the phenol core.

The Epoxigruppen is preferably Epoxialkylreste, which <, e.g. over; RTI ID=3.3> Ether bridges, < /RTI> if necessary in addition, in other way are connected with the bisphenols and/or novolacs.

This Epoxialkylgruppen points e.g. 2 - 5, preferably 3 - 4 Carbons, usually straight-chain up. Particularly prefered is such Epoxi bisphenois and/or Epoxi novolacs, is terminal with which then the Epoxigruppe. Als sehr gut geeignete Beispiele hiorfür werden Bispropoxi-Bisphenole und Monopropoxi-Novolake mit endständigen Epoxigruppen genannt, wobei die Propoxigruppe (eweils mit dem Kern direkt über eine < RIT II D= 4.1 > Achtersausersoftbrücke-(RTD) is connected.

The Epoxiacrylatharze contained in the inventive sticking, detention and sealants < preferably a molecular weight of under 1500, in particular under 1200, although higher molecular weights, e.g. until 2000 or; RTI ID=4.2> darbber < RTI > excluded are not. The reactive resins contained in the inventive means can by in fig. 1 and fig. 2

represented prototypes explained become. Their production can for example in from fig. 3 apparent way take place.

Particularly preserved has itself from the type of bisphenol of derived reactive resins of the kind in speech, which < also in the mixture also of, RTI ID-4.3 > Novolak Typenc /RTI> derived reactive resins that it local skin contained to here can, whereby the content can amount to on from Novolak types derived reactive resins preferably up to 60% and in particular up to 90%, related to the overall weight from the two derementioned reactive resins.

As hardeners for the radical hardenable reactive resins initiators know such as Diarylperoxide, e.g., Dibenzoylperoxid (BP) or to (4-Chiohenzoyl) peroxide (CLBP), Kotonperoxide, e.g., Methylethylkotenperoxid (MEKP) or Cyclohexanonperoxid (CHP), alkyl by esters, e.g., t-Butyl-perbenzoat (TBPB) and such contained to be. They are in usual quantities. e.g., between 0.5 to 5 Gew. - %; related to the total weight, contained.

The initiators can become phlegmatisiert in pure form (e.g., IBPB) or (e.g., with yppsum, chalk, Phthalat, chlorinated paraffin, water) as powders, emulsion or paste used. Further aggregates can be such as fillers contained. When strengthening fillers serve e.g. quartz, glass, corundum, porcelaim, stoneware, Schwerspat, chalk, whereby fillers in the form of Sanden, flours or special < RTI ID=5.1> Formkdrpern</ri>

The bisphenol A carylates and - methacrylate as well as the bisphenol Facrylates and - methacrylate have for example molecular weights within the range of 450 to screedly 800, while the Novolakacrylate and - methacrylate, independing on the number of the phenol cores, e.g., three to eight of the Ausgangsnovolakepoxids, molecular weights of for example within the range of 780 to 2000 to have been able of

< RTI ID-5.2> Reaktivverdtinner / RTID are not compellingly required. They can be used however simultaneous for the
engagement of the desired viscosity. As < RTI ID-5.3> Reaktiverddinner / RTID can then the for this known
connections, for example monostyrene, diviny/benzene, allyl ester more-basic acids, methacrylic acid ethyl esters,
Methacrylsaure i more propylester, Methacrylsaure i more butylester and such a thing be used.

The resin solutions can be before-accelerated with accelerators, how them < for; RTI ID=5.4> cold-hardening, < /RTI> insatlated polyester resins are usual. As such accelerators can become for example Dimethylanilin, Ditthylanilin, Dimethylanilin, Cobaltodoat, Cobaltnaphthenat as well as cobalt/amine accelerator mixtures user.

The advantages of the inventive means lie lirst in the fact that, in particular good ages and < the positive characteristics of the more-full of seeds system; RTI 10-6.15 . Witterrungsbesten < /RTI s into the means introduced become digkeit. In addition it comes that become positively affected by the content of hydroxyl groups the adhesion properties. It becomes a favourable weathing of glass, guartz, sand, rock and thus good detention characteristic achievance.

Contrary to cold-hardening Epoxiden, whose application borders are because of their temperature sensitivity with + 50C, the curing of the inventive radical hardening means with peroxides can be accomplished as with the unsaturated polyesters. Even with temperatures of for example - 100C is still another quick cure possible.

A further advantage lies in the fact that given regarding a smaller volume contraction at ester groups, sessinal opposite insatiated polyseter resins, an high resistance is against alkaline saponification. Particular advantage lies however in it that the double bonds, in particular if they are terminal are particularly reactive there sterisch not hindered. Contrary to the double bonds with insatiated polyester resins, sterisch prevented because of their usual arrangement within the polyester chain, the inventive means react both to quick and practically quantitatively with its double bonds. Due to the prefered situation of the reactive group in the inventive means they are also capable therefore, without additive of < RTI ID=6.2.> Reaktiveradionerer. (RTII to dou't-hard or other comonomers.

With use of the inventive means one receives a cured product, which opposite cured, insatiated polyseter resine sesential is tough-more elastic due to the cross-inking possibilities and a higher tear stretch exhibits. Due to the high RTI ID=7.1> Relsidehnung

 ID=7.1> Relsidehnung
 /RTI> the means are< in the situation, during; RTI ID=7.2> Hortunggverlauts
 /RTI> to take

Particularly prefered < itself mixtures the subsequent Mi; RTI ID=7.3> schungsverhältnisse, < /RTI> related to the entire mixture, but without hardeners, proved:

Reactive resin 9 - 28 Gew. - < %; RTI [D=7.4 > Reaktivverdünner</ri>
| RTI | Section | RTI | Reactive residual | RTI | Reactive residual | RTI | Reactive residual | RTI | RTI | REACTIVE residual | RTI | RTI | REACTIVE residual | RTI |

Such Phlegmatisierungsmittel is in subordinated quantities, for example within the range of 0,5 to 3.5 Gew. - %, related to the entire mixture, contained, just as accelerator, those, if they are present, usually within the range of 0,005 to 2.0 Gew. - are present %, related to the entire mixture.

The use of the inventive means is particularly favourable to the fixture of dowels and mooring rings in Boreholes. Reaction resis and hardeness with the desired in each case constituents can become separate or into the Borehole introduced common immediately after mixing. Usually however reactive resin and hardeners in two chambers are separate contained for this purpose in a unit, e.g., in a cartridge. The cartridge is shen destroyed after bringing into the Borehole by inserting and turning of the dowel and/or the mooring ring, whereby the wall material can serve the cartridge as filler, and then the filter portions to add is. In addition, it is possible to use systems in those the one component, e.g., the hardener, microencepsulesties is. When using the mooring ring the wall material of the microcapsules

The high elongation of break that inventive used means an ensured to a large extent unstressed uptake of the Volumenkontraktion arising during the curing, With burden of the armature an extension of the moving ring dependent of the borehole depth occurs. For inventive tough reading tables the means secure a more favorable load distribution essential opposite known shear connectors on basis of unsaturated polyesters on the entire group range and in particular the degradation at the surface of the photograph material, for the example concrete, arising voltage peak-whole general lead the inventive means, which a mixture of acrylic acid modified bisphenol connections and acrylic acid modified novolac connections contained, to products with higher thermostability, which in particular with the use in the shear connector range, to the fixture of dowels and mooring rings in Boreholes of importance is. This results from from < RTI ID=8.1> Molekillstruktur< /RTI> resulting higher degree of crosslinking. This higher temperature resistance is simultaneous with a higher chemical resistance links a further advantage is the high weathering resistance and the small water uptake also with long-term contact with water.

The inventive means know known reaction resins, like insatiated polyester resins, Methylmethacrylatharze or other known epoxy resins in quantities in the sense of a modification of the characteristics contained, subordinated if necessary

The epoxy bisphenols and/or - novolacs can e.g. become in usual manner by reaction of the bisphenol and/or novolac with epichlorohydrin manufactured (see. e.g. H.G. Elias, macromolecules, Basel, Heidelberg 1972, S. 707 - 709).

In the following embodiments the invention is more near explained.

Example 1 bisphenol A methacrylat 17,90 < RTI ID=9.1> Gew. - E< /RTI> (Mg: 492-775) Novolakmethacrylat 4,30 Gew. - < %; RTI ID=9.2> (Mg; < /RTI> 780-1200) Monostyrene < RTI ID=9.3> 12.20 Gew. - E< /RTI> Divinylbenzene 1,00 Gew. - % silica sand 0.1? 0,25 mm of 30,00 Gew. - % silica sand 0.3? 0.65 < RTI ID=9.4> min< /RTI> 30,10 Gew. - % Dimethyl p toluidin 0,05 Gew. - % Dibenzoylperoxid 4,45 Gew. - % (50% industrial union < in chlorinated paraffin); RTI ID=9.5> 100,00 Gew. - %< /RTI> By mixing one keeps a mass < RTI ID=9.6> putty fähiger< /RTI> Consistency.

Gel time: 3 - 3.5 minutes of curing time: 35 minutes of strength; Mooring ring; M 12 setting depth; 110 mm of borehole diameters; 14 mm concrete; B 35, Istfestigkeit 48N/mm2 failure value; 74 kN eispiel 2 Dimension M 12) Aussenpatrone Glass weight 5.6 g

Glass outside diameter 10,75 mm

Glass wall thickness 0.55 mm

Cartridge length 100 mm

bisphenol A methacrylat (65 < RTI ID=10.1> tig< /RTI> in monostyrene, amine-accelerated) 4.3 g

Quartz (particle size 1.5 - 2.0 mm) 6.6 q) interior cartridge

Glass weight 1.2 g Glass outside diameter 6.30 mm

Glaswandstärke 0.65 mm

Cartridge length 80 mm

Dibenzoylperoxid (20% in gypsum) 0.8 g eispiel 3) outer cartridge

Glass weight 5.8 g

Glass outside diameter 9,20 mm

Glass wall thickness 0,45 mm

Cartridge length 90 mm bisphenol A acrylate (55 < RTI ID=10.2> tig< /RTI> in monostyrene,

40 parts, Divinviben zol 5 parts, beschleu nigt with 0,15%

Dimethyl-p-toluidin) 3,8 g b) Innenpatrone

Glasgewicht 2,2 g

Glass outside diameter 10.75 mm

Glass wall thickness 0,45 mm

Cartridge length 90 mm

Quartz (1.2 - 1.8 mm) 5.4 q

Dibenzoylperoxid (50% industrial union in Phtalat) 0.8 q example 4 A) outer cartridge Glass weight 5.8 g

Glass outside diameter 10.75 mm

Glass wall thickness 0,45 mm

Cartridge length 110 mm

bisphenol A methacrylat (80% industrial union in methacrylic acid ester, before-accelerated also Amine) 4.2 g b) interior cartridge

Glass weight 2.2 g

Glass outside diameter 9,20 mm Glass wall thickness 0.45 mm

Cartridge length 90 mm

Quartz (0.04 - 0.15 mm) 0.4 g

Quartz (1.2 - 1.8 mm) 5.0 g

Dibenzoylperoxid (50% industrial union in chalk) 0.5 g Example â) Outer cartridge

Glass weight 5.6 g Glass outside diameter 10.75 mm

Glass wall thickness 0,45 mm

Cartridge length 100 mm

bisphenol A methacrylat

MG: 492 - 775 (75 < RTI ID=12.1> tig< /RTI> in monostyrene) 3.0 g Novolakmethacrylat,

Mg: 780 - 1200 (60% industrial union in monostyrene) 1.2 g

Cobaltoctoat 0.001 g

Dimethylanilin 0.004 g b) interior cartridge Glass weight 0.7 g

Glass outside diameter 43 mm

Glass wall thickness 0.65 mm

Patronenlänge 80 mm

Methylethylketonperoxid 0.35 g in the images 1 and 2 are in the inventive means contained reaction products in exemplary form explained. It concerns here preferable embodiments. The expiration to the formation of the reaction products is in image 3 in exemplary form schematically shown.